

* * * * * Welcome to STN International * * * * *

NEWS 1 Web Page URLs for STN Seminar Schedule - N. America
 NEWS 2 "Ask CAS" for self-help around the clock
 NEWS 3 May 12 EXTEND option available in structure searching
 NEWS 4 May 12 Polymer links for the POLYLINK command completed in REGISTRY
 NEWS 5 May 27 New UPM (Update Code Maximum) field for more efficient patent
 SDIs in CPlus
 NEWS 6 May 27 CPlus super roles and document types searchable in REGISTRY
 NEWS 7 Jun 22 STN Patent Forums to be held July 19-22, 2004
 NEWS 8 Jun 28 Additional enzyme-catalyzed reactions added to CASREACT
 NEWS 9 Jun 28 ANTE, AQUALINE, BIOENG, CIVILENG, ENVIROENG, MECHENG,
 and WATER from CSA now available on STN(R)

NEWS EXPRESS MARCH 31 CURRENT WINDOWS VERSION IS V7.00A, CURRENT
 MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
 AND CURRENT DISCOVER FILE IS DATED 26 APRIL 2004

NEWS HOURS STN Operating Hours Plus Help Desk Availability
 NEWS INTER General Internet Information
 NEWS LOGIN Welcome Banner and News Items
 NEWS PHONE Direct Dial and Telecommunication Network Access to STN
 NEWS WWW CAS World Wide Web Site (general information)

Enter NEWS followed by the item number or name to see news on that
 specific topic.

All use of STN is subject to the provisions of the STN Customer
 agreement. Please note that this agreement limits use to scientific
 research. Use for software development or design or implementation
 of commercial gateways or other similar uses is prohibited and may
 result in loss of user privileges and other penalties.

* * * * * STN Columbus * * * * *

FILE 'HOME' ENTERED AT 15:26:37 ON 09 JUL 2004

=> file reg		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	0.21	0.21

FILE 'REGISTRY' ENTERED AT 15:26:43 ON 09 JUL 2004
 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
 PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
 COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file
 provided by InfoChem.

STRUCTURE FILE UPDATES: 8 JUL 2004 HIGHEST RN 706430-72-0
 DICTIONARY FILE UPDATES: 8 JUL 2004 HIGHEST RN 706430-72-0

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

Please note that search-term pricing does apply when

conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at:

<http://www.cas.org/ONLINE/DBSS/registryss.html>

=> e 310427-67-9/rn

E1	1	310427-65-7/RN
E2	1	310427-66-8/RN
E3	1 -->	310427-67-9/RN
E4	1	310427-68-0/RN
E5	1	310427-69-1/RN
E6	1	310427-70-4/RN
E7	1	310427-71-5/RN
E8	1	310427-72-6/RN
E9	1	310427-73-7/RN
E10	1	310427-74-8/RN
E11	1	310427-75-9/RN
E12	1	310427-76-0/RN

=> s e3

L1 1 310427-67-9/RN

=> d l1

L1 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN

RN 310427-67-9 REGISTRY

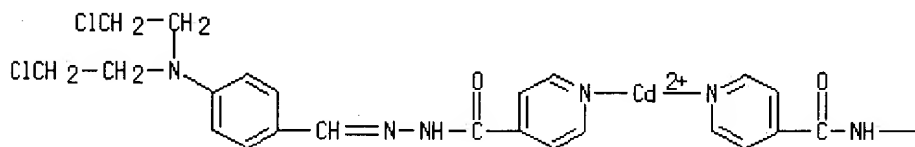
CN Cadmium(2+), bis[(4-pyridinecarboxylic acid-κN1)
[[4-[bis(2-chloroethyl)amino]phenyl]methylene]hydrazide]- (9CI) (CA INDEX
NAME)

MF C34 H36 Cd Cl4 N8 O2

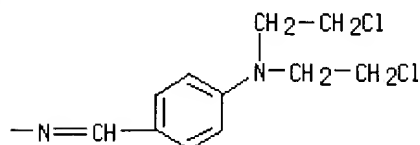
CI CCS, COM

SR CA

PAGE 1-A



PAGE 1-B



=> e 310427-65-7/rn

E1	1	310427-63-5/RN
E2	1	310427-64-6/RN
E3	1 -->	310427-65-7/RN
E4	1	310427-66-8/RN
E5	1	310427-67-9/RN

E6 1 310427-68-0/RN
 E7 1 310427-69-1/RN
 E8 1 310427-70-4/RN
 E9 1 310427-71-5/RN
 E10 1 310427-72-6/RN
 E11 1 310427-73-7/RN
 E12 1 310427-74-8/RN

=> s e3

L2 1 310427-65-7/RN

=> d 12

L2 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN

RN 310427-65-7 REGISTRY

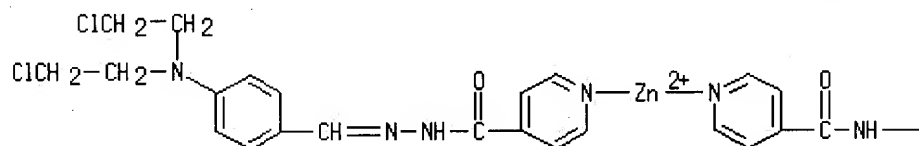
CN Zinc(2+), bis[(4-pyridinecarboxylic acid-κN1) [[4-[bis(2-chloroethyl)amino]phenyl]methylene]hydrazide] - (9CI) (CA INDEX NAME)

MF C34 H36 Cl4 N8 O2 Zn

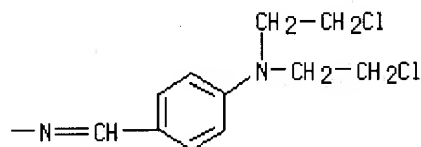
CI CCS, COM

SR CA

PAGE 1-A



PAGE 1-B



=> e 10325-94-7/rn

E1 1 10325-88-9/RN
 E2 1 10325-89-0/RN
 E3 1 --> 10325-94-7/RN
 E4 1 103250-00-6/RN
 E5 1 103250-01-7/RN
 E6 1 103250-02-8/RN
 E7 1 103250-03-9/RN
 E8 1 103250-04-0/RN
 E9 1 103250-05-1/RN
 E10 1 103250-06-2/RN
 E11 1 103250-07-3/RN
 E12 1 103250-08-4/RN

=> s e3

L3 1 10325-94-7/RN

=> d 13

L3 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN

RN 10325-94-7 REGISTRY

CN Nitric acid, cadmium salt (8CI, 9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Cadmium nitrate (7CI)

OTHER NAMES:

CN Cadmium dinitrate

CN Cadmium nitrate (Cd(NO3)2)

CN Cadmium(II) nitrate

DR 14177-24-3

MF Cd . 2 H N O3

CI COM

LC STN Files: AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS, BIOTECHNO,
CA, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST, CIN, CSCHM,
CSNB, DETHERM*, EMBASE, GMELIN*, HSDB*, IFICDB, IFIPAT, IFIUDB, MEDLINE,
MRCK*, MSDS-OHS, NIOSHTIC, PDLCOM*, PIRA, PROMT, RTECS*, TOXCENTER,
USPAT2, USPATFULL

(*File contains numerically searchable property data)

Other Sources: DSL**, EINECS**, TSCA**

(**Enter CHEMLIST File for up-to-date regulatory information)

DT.CA Caplus document type: Conference; Dissertation; Journal; Patent; Report

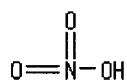
RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study);
MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC
(Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses);
NORL (No role in record)

RLD.P Roles for non-specific derivatives from patents: PROC (Process); RACT
(Reactant or reagent); USES (Uses)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological
study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU
(Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT
(Reactant or reagent); USES (Uses); NORL (No role in record)

RLD.NP Roles for non-specific derivatives from non-patents: BIOL (Biological
study); PREP (Preparation); PRP (Properties); RACT (Reactant or
reagent); USES (Uses)

CRN (7697-37-2)



1/2 Cd

1477 REFERENCES IN FILE CA (1907 TO DATE)

19 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

1480 REFERENCES IN FILE CAPLUS (1907 TO DATE)

1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=>

* * * * * RECONNECTED TO STN INTERNATIONAL * * * * *

SESSION RESUMED IN FILE 'REGISTRY' AT 15:39:18 ON 09 JUL 2004

FILE 'REGISTRY' ENTERED AT 15:39:18 ON 09 JUL 2004

COPYRIGHT (C) 2004 American Chemical Society (ACS)

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	14.13	14.34

=> file casreact

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	14.97	15.18

FILE 'CASREACT' ENTERED AT 15:40:30 ON 09 JUL 2004
 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT
 COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications.

FILE CONTENT:1840 - 4 Jul 2004 VOL 141 ISS 1

 *
 * CASREACT now has more than 8 million reactions *
 *

Some CASREACT records are derived from the ZIC/VINITI database (1974-1991) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=>

L4 STRUCTURE UPLOADED

=> d 14

L4 HAS NO ANSWERS

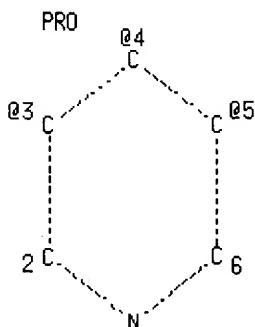
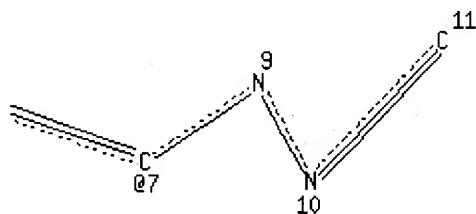
L4 STR

0 24 S 25

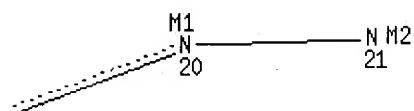
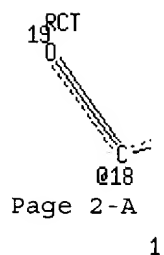
PRO

80

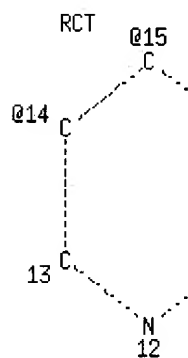
Page 1-A



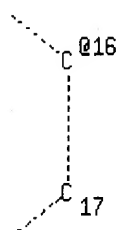
Page 1-B



Page 2-B



Page 3-A



Page 3-B

VAR G1=24/25

VPA 7-3/4/5 S

VPA 18-14/15/16 S

NODE ATTRIBUTES:

RCT
22



HCOUNT IS M1 AT 20
 HCOUNT IS M2 AT 21
 NSPEC IS R AT 1
 NSPEC IS R AT 2
 NSPEC IS R AT 3
 NSPEC IS R AT 4
 NSPEC IS R AT 5
 NSPEC IS R AT 6
 NSPEC IS C AT 7
 NSPEC IS C AT 8
 NSPEC IS C AT 9
 NSPEC IS C AT 10
 NSPEC IS RC AT 11
 NSPEC IS R AT 12
 NSPEC IS R AT 13
 NSPEC IS R AT 14
 NSPEC IS R AT 15
 NSPEC IS R AT 16
 NSPEC IS R AT 17
 NSPEC IS C AT 18
 NSPEC IS C AT 19
 NSPEC IS C AT 20
 NSPEC IS C AT 21
 NSPEC IS C AT 22
 NSPEC IS RC AT 23
 DEFAULT MLEVEL IS ATOM
 MLEVEL IS CLASS AT 7 8 9 10 11 18 19 20 21 23 24 25
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

=> s 14

SAMPLE SEARCH INITIATED 15:45:45 FILE 'CASREACT'

SCREENING COMPLETE - 147 REACTIONS TO VERIFY FROM 27 DOCUMENTS

100.0% DONE 147 VERIFIED 53 HIT RXNS 11 DOCS
 SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
 BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 2213 TO 3667
 PROJECTED ANSWERS: 22 TO 418

L5 11 SEA SSS SAM L4 (53 REACTIONS)

=> s 14 full

THE ESTIMATED SEARCH COST FOR FILE 'CASREACT' IS 102.30 U.S. DOLLARS

DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 15:45:49 FILE 'CASREACT'

SCREENING COMPLETE - 3426 REACTIONS TO VERIFY FROM 643 DOCUMENTS

100.0% DONE 3426 VERIFIED 618 HIT RXNS 187 DOCS
 SEARCH TIME: 00.00.01

L6 187 SEA SSS FUL L4 (618 REACTIONS)

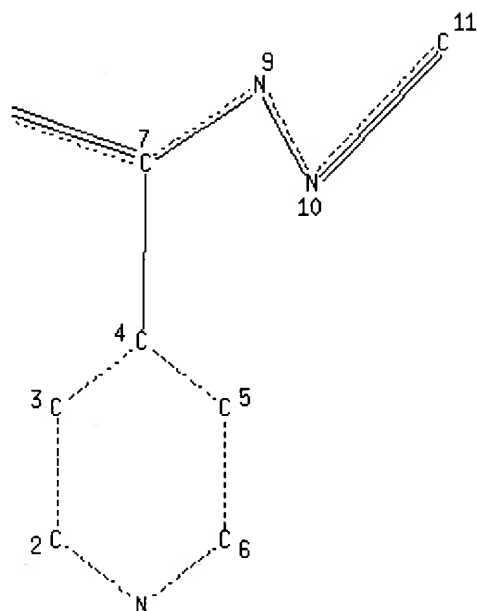
=>
L7 STRUCTURE UPLOADED

=> d 17
L7 HAS NO ANSWERS
L7 STR
0 24 S 25

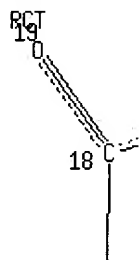
PRO

8 0

Page 1-A

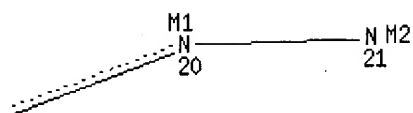


Page 1-B

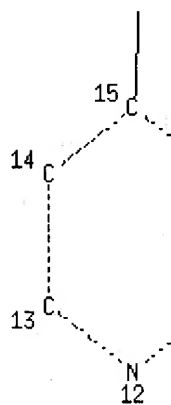


Page 2-A

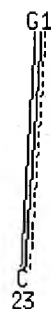
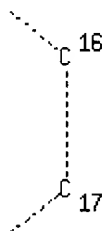
1

RGT
22

Page 2-B



Page 3-A



Page 3-B

VAR G1=24/25

NODE ATTRIBUTES:

HCOUNT	IS	M1	AT	20
HCOUNT	IS	M2	AT	21
NSPEC	IS	R	AT	1
NSPEC	IS	R	AT	2
NSPEC	IS	R	AT	3
NSPEC	IS	R	AT	4
NSPEC	IS	R	AT	5

NSPEC IS R AT 6
 NSPEC IS C AT 7
 NSPEC IS C AT 8
 NSPEC IS C AT 9
 NSPEC IS C AT 10
 NSPEC IS RC AT 11
 NSPEC IS R AT 12
 NSPEC IS R AT 13
 NSPEC IS R AT 14
 NSPEC IS R AT 15
 NSPEC IS R AT 16
 NSPEC IS R AT 17
 NSPEC IS C AT 18
 NSPEC IS C AT 19
 NSPEC IS C AT 20
 NSPEC IS C AT 21
 NSPEC IS C AT 22
 NSPEC IS RC AT 23
 DEFAULT MLEVEL IS ATOM
 MLEVEL IS CLASS AT 7 8 9 10 11 18 19 20 21 23 24 25
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

=> s 17

SAMPLE SEARCH INITIATED 15:47:23 FILE 'CASREACT'

SCREENING COMPLETE - 50 REACTIONS TO VERIFY FROM 13 DOCUMENTS

100.0% DONE 50 VERIFIED 24 HIT RXNS 10 DOCS
 SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
 BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 576 TO 1424
 PROJECTED ANSWERS: 11 TO 389

L8 10 SEA SSS SAM L7 (24 REACTIONS)

=> s 17 full

THE ESTIMATED SEARCH COST FOR FILE 'CASREACT' IS 102.30 U.S. DOLLARS

DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 15:47:28 FILE 'CASREACT'

SCREENING COMPLETE - 957 REACTIONS TO VERIFY FROM 233 DOCUMENTS

100.0% DONE 957 VERIFIED 443 HIT RXNS 157 DOCS
 SEARCH TIME: 00.00.01

L9 157 SEA SSS FUL L7 (443 REACTIONS)

=> file reg

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

209.64

224.82

FILE 'REGISTRY' ENTERED AT 15:47:49 ON 09 JUL 2004

USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
 PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
 COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file
 provided by InfoChem.

STRUCTURE FILE UPDATES: 8 JUL 2004 HIGHEST RN 706430-72-0
 DICTIONARY FILE UPDATES: 8 JUL 2004 HIGHEST RN 706430-72-0

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

Please note that search-term pricing does apply when
 conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more
 information enter HELP PROP at an arrow prompt in the file or refer
 to the file summary sheet on the web at:
<http://www.cas.org/ONLINE/DBSS/registryss.html>

=> e absolute ethanol/cn

E1	1	ABSOLAC SP 200/CN
E2	1	ABSOLUTE/CN
E3	0 -->	ABSOLUTE ETHANOL/CN
E4	1	ABSON/CN
E5	1	ABSON 69163/CN
E6	1	ABSON 820X14/CN
E7	1	ABSON 820X17/CN
E8	1	ABSON 821/CN
E9	1	ABSON 89110/CN
E10	1	ABSON 89120/CN
E11	1	ABSON 89131/CN
E12	1	ABSON 89140/CN

=> file casreact

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	0.42	225.24

FILE 'CASREACT' ENTERED AT 15:48:05 ON 09 JUL 2004
 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT
 COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is
 held by the publishers listed in the PUBLISHER (PB) field (available
 for records published or updated in Chemical Abstracts after December
 26, 1996), unless otherwise indicated in the original publications.

FILE CONTENT:1840 - 4 Jul 2004 VOL 141 ISS 1

```
*****
*
*   CASREACT now has more than 8 million reactions
*
*****
```

Some CASREACT records are derived from the ZIC/VINITI database (1974-1991)
 provided by InfoChem, INPI data prior to 1986, and Biotransformations

database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

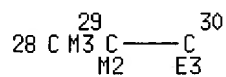
=>

L10 STRUCTURE UPLOADED

=> d 110

L10 HAS NO ANSWERS

L10 STR

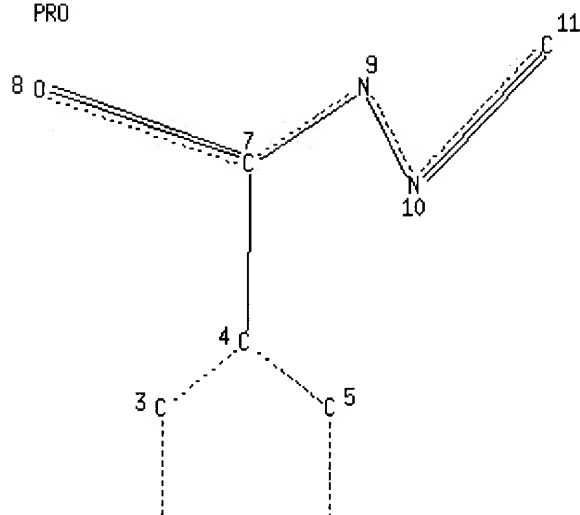


0 26 S 27

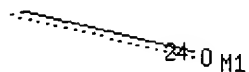
RRT
25 G2

Page 1-C

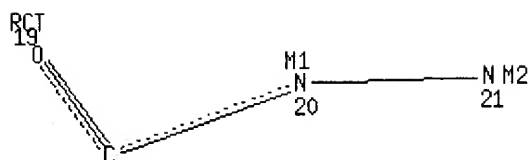
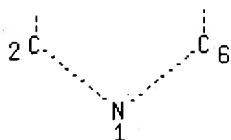
PRO



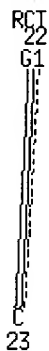
Page 1-D



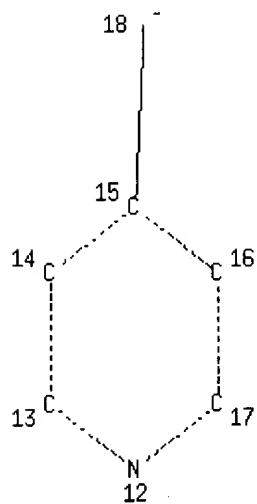
Page 2-C



Page 2-D



Page 3-A



Page 3-D

VAR G1=26/27

VAR G2=28/29

NODE ATTRIBUTES:

HCOUNT IS M1 AT 20

HCOUNT IS M2 AT 21
 HCOUNT IS M1 AT 24
 HCOUNT IS M3 AT 28
 HCOUNT IS M2 AT 29
 HCOUNT IS E3 AT 30
 NSPEC IS R AT 1
 NSPEC IS R AT 2
 NSPEC IS R AT 3
 NSPEC IS R AT 4
 NSPEC IS R AT 5
 NSPEC IS R AT 6
 NSPEC IS C AT 7
 NSPEC IS C AT 8
 NSPEC IS C AT 9
 NSPEC IS C AT 10
 NSPEC IS RC AT 11
 NSPEC IS R AT 12
 NSPEC IS R AT 13
 NSPEC IS R AT 14
 NSPEC IS R AT 15
 NSPEC IS R AT 16
 NSPEC IS R AT 17
 NSPEC IS C AT 18
 NSPEC IS C AT 19
 NSPEC IS C AT 20
 NSPEC IS C AT 21
 NSPEC IS C AT 22
 NSPEC IS RC AT 23
 NSPEC IS C AT 24
 NSPEC IS C AT 25
 DEFAULT MLEVEL IS ATOM
 MLEVEL IS CLASS AT 7 8 9 10 11 18 19 20 21 23 24 26 27 28 29 30
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 30

STEREO ATTRIBUTES: NONE

=> s 110

SAMPLE SEARCH INITIATED 15:50:55 FILE 'CASREACT'

SCREENING COMPLETE - 50 REACTIONS TO VERIFY FROM 13 DOCUMENTS

100.0% DONE 50 VERIFIED 6 HIT RXNS 1 DOCS
 SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 576 TO 1424

PROJECTED ANSWERS: 1 TO 79

L11 1 SEA SSS SAM L10 (6 REACTIONS)

=> s 111 full

THE ESTIMATED SEARCH COST FOR FILE 'CASREACT' IS 102.30 U.S. DOLLARS

DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 15:51:02 FILE 'CASREACT'

SCREENING COMPLETE - 956 REACTIONS TO VERIFY FROM 232 DOCUMENTS

100.0% DONE 956 VERIFIED 9 HIT RXNS 3 DOCS
 SEARCH TIME: 00.00.01

L12 3 SEA SSS FUL L10 (9 REACTIONS)

=> d l12, crd rxn ibib abs, 1-3
 'RXN' IS NOT A VALID FORMAT FOR FILE 'CASREACT'

The following are valid formats:

ABS ----- GI and AB
 ALL ----- BIB, AB, IND, RE, Single-step Reactions
 APPS ----- AI, PRAI
 BIB ----- AN, plus Bibliographic Data
 CAN ----- List of CA abstract numbers without answer numbers
 CBIB ----- AN, plus Compressed Bibliographic Data
 DALL ----- ALL, delimited (end of each field identified)
 IABS ----- ABS, indented with text labels
 IALL ----- ALL, indented with text labels
 IBIB ----- BIB, indented with text labels
 IND ----- Indexing data
 IPC ----- International Patent Classifications
 ISTD ----- STD, indented with text labels
 OBIB ----- AN, plus Bibliographic Data (original)
 OIBIB ----- OBIB, indented with text labels

SBIB ----- BIB, no citations
 SIBIB ----- IBIB, no citations

MAX ----- Same as ALL
 PATS ----- PI, SO
 SCAN ----- TI and FCRD (random display, no answer number. SCAN
 must be entered on the same line as DISPLAY, e.g.,
 D SCAN.)

SSRX ----- Single-Step Reactions (Map, Diagram, and Summary for
 all single-step reactions)
 STD ----- BIB, IPC, and NCL

CRD ----- Compact Display of All Hit Reactions
 CRDREF ----- Compact Reaction Display and SO, PY for Reference
 FHIT ----- Reaction Map, Diagram, and Summary for first
 hit reaction
 FHITCBIB --- FHIT, AN plus CBIB
 FCRD ----- First hit in Compact Reaction Display (CRD) format
 FCRDREF ----- First hit in Compact Reaction Display (CRD) format with
 CA reference information (SO, PY). (Default)
 FPATH ----- PATH, plus Reaction Summary for the "long path"
 FSPATH ----- SPATH, plus Reaction Summary for the "short path"
 HIT ----- Reaction Map, Reaction Diagram, and Reaction
 Summary for all hit reactions and fields containing
 hit terms
 OCC ----- All hit fields and the number of occurrences of the
 hit terms in each field. Includes total number of
 HIT, PATH, SPATH reactions. Labels reactions that have
 incomplete verifications.
 PATH ----- Reaction Map and Reaction Diagram for the "long
 path". Displays all hit reactions, except those
 whose steps are totally included within another hit
 reaction which is displayed

RX ----- Hit Reactions (Map, Diagram, Summary for all hit reactions)
 RXG ----- Hit Reaction Graphics (Map and Diagram for all hit reactions)
 RXL ----- Hit Reaction Long (Map, Diagram, Summary for all hit reactions)
 RXS ----- Hit Reaction Summaries (Map and Summary for all hit reactions)
 SPATH ----- Reaction Map and Reaction Diagram for the "short
 path". Displays all single step reactions which
 contain a hit substance. Also displays those
 multistep reactions that have a hit substance in both
 the first and last steps of the reaction, except for
 those hit reactions whose steps are totally included
 within another hit reaction which is displayed

To display a particular field or fields, enter the display field
 codes. For a list of the display field codes, enter HELP DFIELDS
 at an arrow prompt (=>). Examples of combinations include: D TI;
 D BIB RX; D TI, AU, FCRD. The information is displayed in the same order
 as the specification. All of the formats, except CRD, CRDREF, FHIT, PATH,
 FPATH, SPATH, FSPATH, FCRD, FCRDREF, HIT, RX, RXG, RXS, SCAN, and OCC, may
 be used with the DISPLAY command to display the record for a specified
 Accession Number.

ENTER DISPLAY FORMAT (FCRDREF):end

=> d his

(FILE 'HOME' ENTERED AT 15:26:37 ON 09 JUL 2004)

FILE 'REGISTRY' ENTERED AT 15:26:43 ON 09 JUL 2004

E 310427-67-9/RN
 L1 1 S E3
 E 310427-65-7/RN
 L2 1 S E3
 E 10325-94-7/RN
 L3 1 S E3

FILE 'CASREACT' ENTERED AT 15:40:30 ON 09 JUL 2004

L4 STRUCTURE UPLOADED
 L5 11 S L4
 L6 187 S L4 FULL
 L7 STRUCTURE UPLOADED
 L8 10 S L7
 L9 157 S L7 FULL

FILE 'REGISTRY' ENTERED AT 15:47:49 ON 09 JUL 2004
 E ABSOLUTE ETHANOL/CN

FILE 'CASREACT' ENTERED AT 15:48:05 ON 09 JUL 2004

L10 STRUCTURE UPLOADED
 L11 1 S L10
 L12 3 S L11 FULL

=> d l12, ibib abs fhit abs, 1-3

L12 ANSWER 1 OF 3 CASREACT COPYRIGHT 2004 ACS on STN

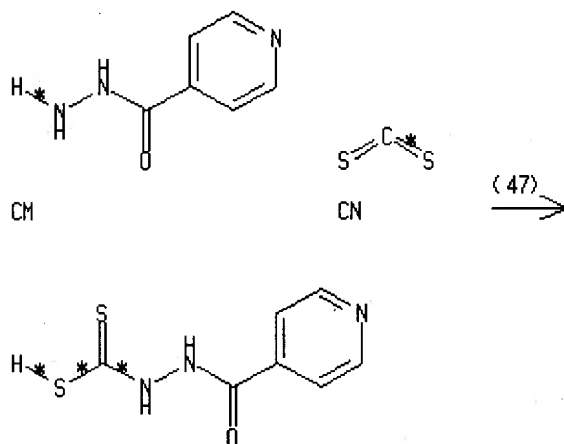
Full Citing
 Text References

ACCESSION NUMBER: 138:39232 CASREACT
 TITLE: Synthesis and biological activity of some
 1,2,4-triazoles and related heterocycles

AUTHOR(S): Udipi, R. H.; Kulkarni, V. M.; Sudheendra; Setty, S.
 Ramachandra; Purushottamachar, P.
 CORPORATE SOURCE: Department of Pharmaceutical Chemistry, V.L. College
 of Pharmacy, Raichur, 584 101, India
 SOURCE: Indian Journal of Heterocyclic Chemistry (2002),
 11(4), 303-308
 CODEN: IJCHEI; ISSN: 0971-1627
 PUBLISHER: Prof. R. S. Varma
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB 4-Amino-2,4-dihydro-5-(4-pyridinyl)-3H-1,2,4-triazole-3-thione and
 5-(4-pyridinyl)-4-[[[(4-pyridinyl)carbonyl]amino]-3H-1,2,4-triazole-3-
 thione were prep'd.; cyclocondensation of the latter with aryl carboxylic
 acids or aryl aldehydes gave 6-aryl-3-(4-pyridinyl)-1,2,4-triazolo[3,4-
 b][1,3,4]thiadiazole derivs. Some of them were found to be potential
 antiinflammatory and antimicrobial agents.

RX(47) OF 143 CM + CN ==> CO...



K

CO

RX(47) RCT CM 54-85-3, CN 75-15-0
 RGT CP 1310-58-3 KOH, CQ 64-17-5 EtOH, BM 7732-18-5 Water
 PRO CO 61019-32-7

AB 4-Amino-2,4-dihydro-5-(4-pyridinyl)-3H-1,2,4-triazole-3-thione and
 5-(4-pyridinyl)-4-[[[(4-pyridinyl)carbonyl]amino]-3H-1,2,4-triazole-3-
 thione were prep'd.; cyclocondensation of the latter with aryl carboxylic
 acids or aryl aldehydes gave 6-aryl-3-(4-pyridinyl)-1,2,4-triazolo[3,4-
 b][1,3,4]thiadiazole derivs. Some of them were found to be potential
 antiinflammatory and antimicrobial agents.

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 2 OF 3 CASREACT COPYRIGHT 2004 ACS on STN

Full Citing
 Text References

ACCESSION NUMBER: 135:131724 CASREACT

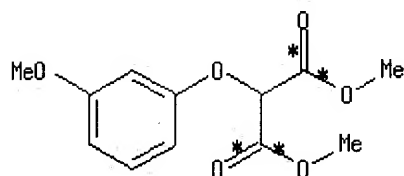
TITLE: Synthesis and structure-activity relationships of

potent and orally active sulfonamide ETB selective antagonists

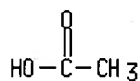
AUTHOR(S): Kanda, Y.; Kawanishi, Y.; Oda, K.; Sakata, T.; Mihara, S.; Asakura, K.; Kanemasa, T.; Ninomiya, M.; Fujimoto, M.; Konoike, T.
 CORPORATE SOURCE: Shionogi Research Laboratories, Shionogi & Co., Ltd., Fukushima-ku, Osaka, 553-0002, Japan
 SOURCE: Bioorganic & Medicinal Chemistry (2001), 9(4), 897-907
 CODEN: BMECEP; ISSN: 0968-0896
 PUBLISHER: Elsevier Science Ltd.
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB The synthesis and structure-activity relationships of a series of N-pyrimidinyl benzenesulfonamides as ETB selective antagonists are described. N-Isoxazolyl benzenesulfonamide (I), previously reported, was selected as a lead compd., and isosteric replacement of the isoxazole ring of I with a pyrimidine ring led to the discovery of a highly potent ETB selective antagonist with oral bioavailability. Modification of the terminal aldehyde group at the 6-position of the pyrimidine ring was investigated, and malonate and acylhydrazone derivs. were equipotent to aldehyde deriv. (II). Compd. II showed ETB antagonistic activity on in vivo evaluation.

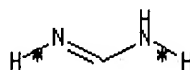
RX(120) OF 135 COMPOSED OF RX(29), RX(2), RX(3), RX(4), RX(30), RX(27)
 RX(120) C + AG + J + M + BT ==> BU



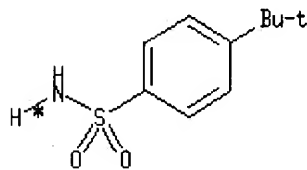
C



AG: CM 1

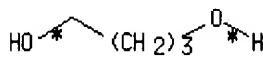


AG: CM 2

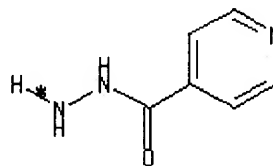


K

J

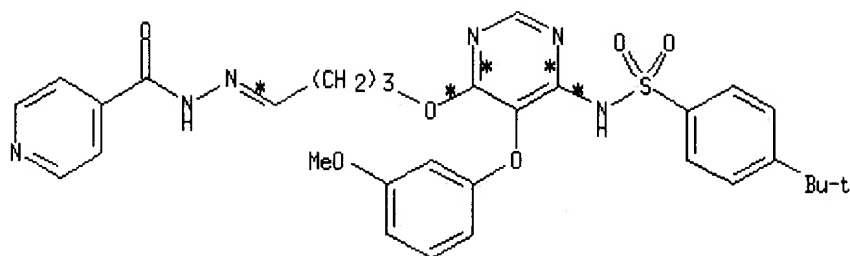


M



BT

6
 STEPS
 →



BU

RX (29) RCT C 189574-46-7, AG 3473-63-0
 RGT D 124-41-4 NaOMe
 PRO F 202288-10-6
 SOL 67-56-1 MeOH

RX (2) RCT F 202288-10-6
 RGT H 10025-87-3 POCl3
 PRO G 150727-28-9
 SOL 10025-87-3 POCl3, 108-75-8 s-Collidine

RX (3) RCT G 150727-28-9, J 146796-98-7
 PRO K 150727-29-0
 SOL 67-68-5 DMSO

RX (4) RCT K 150727-29-0, M 110-63-4
 RGT O 7646-69-7 NaH
 PRO N 202288-11-7
 SOL 110-63-4 HO(CH2)4OH

RX (30) RCT N 202288-11-7
 RGT R 26299-14-9 PCC
 PRO AO 202287-80-7
 SOL 75-09-2 CH2Cl2

RX (27) RCT AO 202287-80-7, BT 54-85-3
 PRO BU 231613-18-6
 SOL 64-17-5 EtOH

AB The synthesis and structure-activity relationships of a series of N-pyrimidinyl benzenesulfonamides as ETB selective antagonists are described. N-Isoxazolyl benzenesulfonamide (I), previously reported, was selected as a lead compd., and isosteric replacement of the isoxazole ring of I with a pyrimidine ring led to the discovery of a highly potent ETB selective antagonist with oral bioavailability. Modification of the terminal aldehyde group at the 6-position of the pyrimidine ring was investigated, and malonate and acylhydrazone derivs. were equipotent to aldehyde deriv. (II). Compd. II showed ETB antagonistic activity on in vivo evaluation.

REFERENCE COUNT: 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 3 OF 3 CASREACT COPYRIGHT 2004 ACS on STN

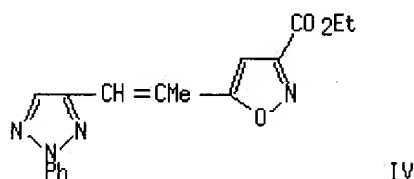
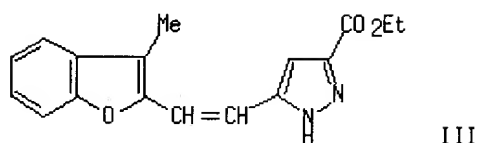
Full Citing
Text References

ACCESSION NUMBER: 104:50817 CASREACT
 TITLE: Synthesis of nitrogenous compounds from
 δ-unsaturated 1,3-dicarbonyl esters:
 trisubstituted pyrazoles of possible antimicrobial and

hypoglycemic activities and hydrazones with antituberculosis activity

AUTHOR(S): Mokhtar, Hassan M.; Wojtanis, J.
 CORPORATE SOURCE: Fac. Sci., Alexandria Univ., Egypt
 SOURCE: Indian Journal of Chemistry, Section B: Organic Chemistry Including Medicinal Chemistry (1985), 24B(2), 188-92
 CODEN: IJSBDB; ISSN: 0376-4699

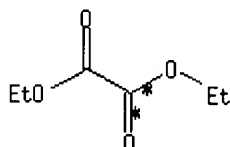
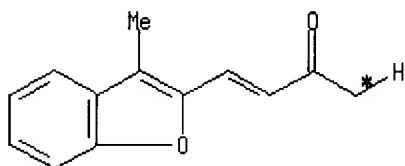
DOCUMENT TYPE: Journal
 LANGUAGE: English
 GI

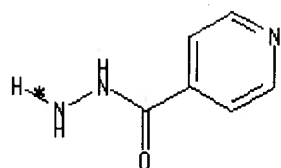


AB R1CH:CRCOCH2COCO2Et (I, R = H, Me, Ph; R1 = 3-methylbenzofuran-2-yl, 2-phenyl-2H-1,2,3-triazol-4-yl) were prepd. by condensing R1CH:CRCOMe (II) with Et oxalate in dry ether. Treating I with hydrazine and arylhydrazines caused cyclization to the corresponding Et substituted pyrazole-3-carboxylates, e.g., III, which were hydrolyzed to acids or converted into acid hydrazides. Condensing I with acylhydrazines gave hydrazones which were cyclized to the corresponding N-acylpyrazoles. Treating I with HONH2 gave 3,5-disubstituted isoxazoles, e.g., IV, whereas using o-phenylenediamine gave hydroxyquinoxalines. Reaction of II with arylhydrazines gave the corresponding hydrazones, which on boiling with EtOH-HCl cyclized to pyrazolines. Treating the latter compds. with excess bromine-water gave the brominated pyrazoles. Condensing II with acylhydrazines gave the corresponding acylhydrazones.

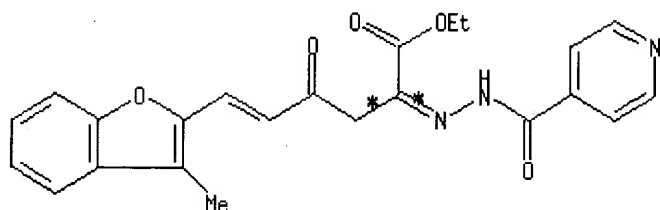
RX(167) OF 375 COMPOSED OF RX(6), RX(85)

RX(167) C + N + DQ ==> DR





DQ

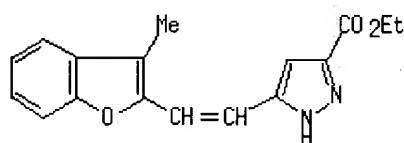
2
STEPS
→

DR

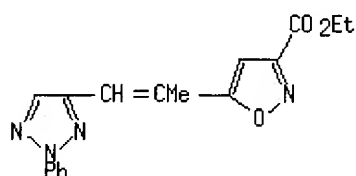
RX(6) RCT C 99410-65-8, N 95-92-1
 RGT P 141-52-6 NaOEt
 PRO O 99410-72-7
 SOL 71-43-2 Benzene, 60-29-7 Et2O

RX(85) RCT DQ 54-85-3, O 99410-72-7
 RGT DB 64-19-7 AcOH
 PRO DR 99411-45-7
 SOL 64-17-5 EtOH

GI



III



IV

AB R1CH:CRCOCH2COCO2Et (I, R = H, Me, Ph; R1 = 3-methylbenzofuran-2-yl, 2-phenyl-2H-1,2,3-triazol-4-yl) were prep'd. by condensing R1CH:CRCOME (II) with Et oxalate in dry ether. Treating I with hydrazine and arylhydrazines caused cyclization to the corresponding Et substituted pyrazole-3-carboxylates, e.g., III, which were hydrolyzed to acids or converted into acid hydrazides. Condensing I with acylhydrazines gave hydrazones which were cyclized to the corresponding N-acylpyrazoles. Treating I with HONH2 gave 3,5-disubstituted isoxazoles, e.g., IV, whereas using o-phenylenediamine gave hydroxyquinoxalines. Reaction of II with arylhydrazines gave the corresponding hydrazones, which on boiling with EtOH-HCl cyclized to pyrazolines. Treating the latter compds. with excess bromine-water gave the brominated pyrazoles. Condensing II with acylhydrazines gave the corresponding acylhydrazones.

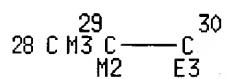
=>

L13 STRUCTURE UPLOADED

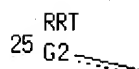
=> d l13

L13 HAS NO ANSWERS

L13 STR

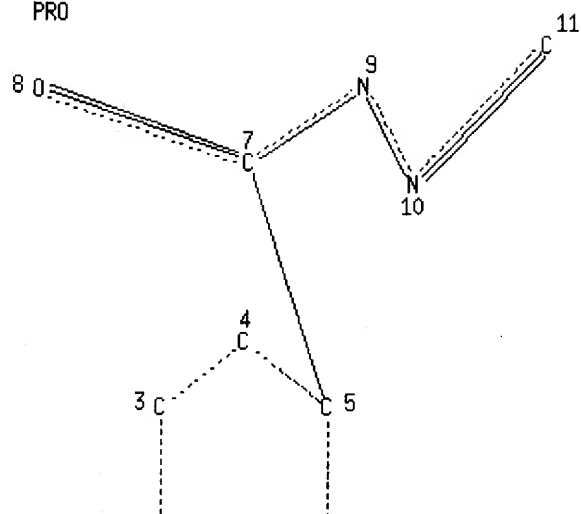


0 26 S 27



Page 1-C

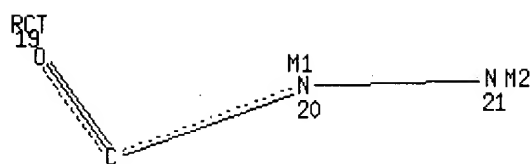
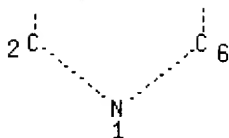
PRO



Page 1-D



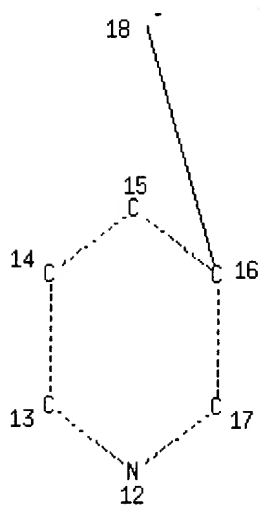
Page 2-C



Page 2-D



Page 3-A



Page 3-D

VAR G1=26/27

VAR G2=28/29

NODE ATTRIBUTES:

HCOUNT IS M1 AT 20

HCOUNT IS M2 AT 21
 HCOUNT IS M1 AT 24
 HCOUNT IS M3 AT 28
 HCOUNT IS M2 AT 29
 HCOUNT IS E3 AT 30
 NSPEC IS R AT 1
 NSPEC IS R AT 2
 NSPEC IS R AT 3
 NSPEC IS R AT 4
 NSPEC IS R AT 5
 NSPEC IS R AT 6
 NSPEC IS C AT 7
 NSPEC IS C AT 8
 NSPEC IS C AT 9
 NSPEC IS C AT 10
 NSPEC IS RC AT 11
 NSPEC IS R AT 12
 NSPEC IS R AT 13
 NSPEC IS R AT 14
 NSPEC IS R AT 15
 NSPEC IS R AT 16
 NSPEC IS R AT 17
 NSPEC IS C AT 18
 NSPEC IS C AT 19
 NSPEC IS C AT 20
 NSPEC IS C AT 21
 NSPEC IS C AT 22
 NSPEC IS RC AT 23
 NSPEC IS C AT 24
 NSPEC IS C AT 25
 DEFAULT MLEVEL IS ATOM
 MLEVEL IS CLASS AT 7 8 9 10 11 18 19 20 21 23 24 26 27 28 29 30
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 30

STEREO ATTRIBUTES: NONE

=> s 113

SAMPLE SEARCH INITIATED 15:55:44 FILE 'CASREACT'

SCREENING COMPLETE - 33 REACTIONS TO VERIFY FROM 7 DOCUMENTS

100.0% DONE 33 VERIFIED 6 HIT RXNS 1 DOCS
 SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 316 TO 1004

PROJECTED ANSWERS: 1 TO 79

L14 1 SEA SSS SAM L13 (6 REACTIONS)

=> s 113 full

THE ESTIMATED SEARCH COST FOR FILE 'CASREACT' IS 102.30 U.S. DOLLARS

DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 15:55:53 FILE 'CASREACT'

SCREENING COMPLETE - 419 REACTIONS TO VERIFY FROM 77 DOCUMENTS

100.0% DONE 419 VERIFIED 6 HIT RXNS
SEARCH TIME: 00.00.01

1 DOCS

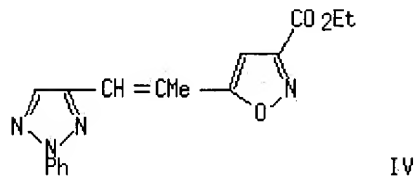
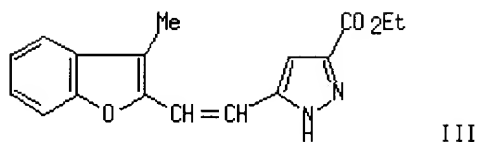
L15 1 SEA SSS FUL L13 (6 REACTIONS)

=> d l15, ibib abs fhit, 1

L15 ANSWER 1 OF 1 CASREACT COPYRIGHT 2004 ACS on STN

Full Text	Citing References
--------------	----------------------

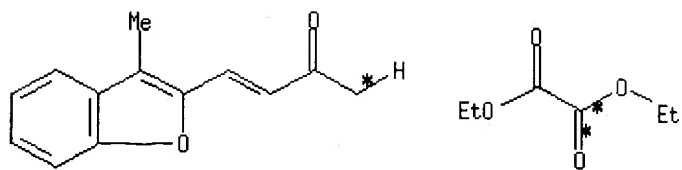
ACCESSION NUMBER:	104:50817 CASREACT
TITLE:	Synthesis of nitrogenous compounds from δ-unsaturated 1,3-dicarbonyl esters: trisubstituted pyrazoles of possible antimicrobial and hypoglycemic activities and hydrazones with antituberculosis activity
AUTHOR(S):	Mokhtar, Hassan M.; Wojtanis, J.
CORPORATE SOURCE:	Fac. Sci., Alexandria Univ., Egypt
SOURCE:	Indian Journal of Chemistry, Section B: Organic Chemistry Including Medicinal Chemistry (1985), 24B(2), 188-92 CODEN: IJSBDB; ISSN: 0376-4699
DOCUMENT TYPE:	Journal
LANGUAGE:	English
GI	



AB R1CH:CRCOCH2COCO2Et (I, R = H, Me, Ph; R1 = 3-methylbenzofuran-2-yl, 2-phenyl-2H-1,2,3-triazol-4-yl) were prepd. by condensing R1CH:CRCOME (II) with Et oxalate in dry ether. Treating I with hydrazine and arylhydrazines caused cyclization to the corresponding Et substituted pyrazole-3-carboxylates, e.g., III, which were hydrolyzed to acids or converted into acid hydrazides. Condensing I with acylhydrazines gave hydrazones which were cyclized to the corresponding N-acylpyrazoles. Treating I with HONH2 gave 3,5-disubstituted isoxazoles, e.g., IV, whereas using o-phenylenediamine gave hydroxyquinoxalines. Reaction of II with arylhydrazines gave the corresponding hydrazones, which on boiling with EtOH-HCl cyclized to pyrazolines. Treating the latter compds. with excess bromine-water gave the brominated pyrazoles. Condensing II with acylhydrazines gave the corresponding acylhydrazones.

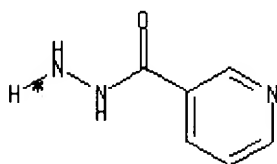
RX(168) OF 375 COMPOSED OF RX(6), RX(86)

RX(168) C + N + DS ==> DT



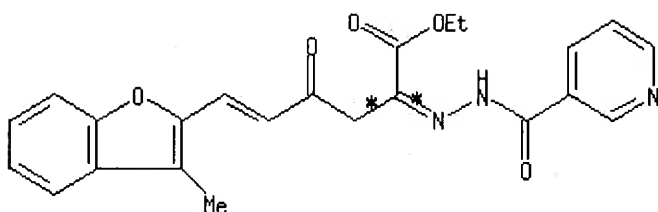
C

N



DS

2
STEPS
→



DT

RX(6) RCT C 99410-65-8, N 95-92-1
 RGT P 141-52-6 NaOEt
 PRO O 99410-72-7
 SOL 71-43-2 Benzene, 60-29-7 Et2O

RX(86) RCT DS 553-53-7, O 99410-72-7
 RGT DB 64-19-7 AcOH
 PRO DT 99411-46-8
 SOL 64-17-5 EtOH

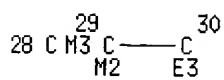
=>

L16 STRUCTURE UPLOADED

=> d 116

L16 HAS NO ANSWERS

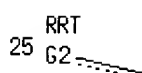
L16 STR



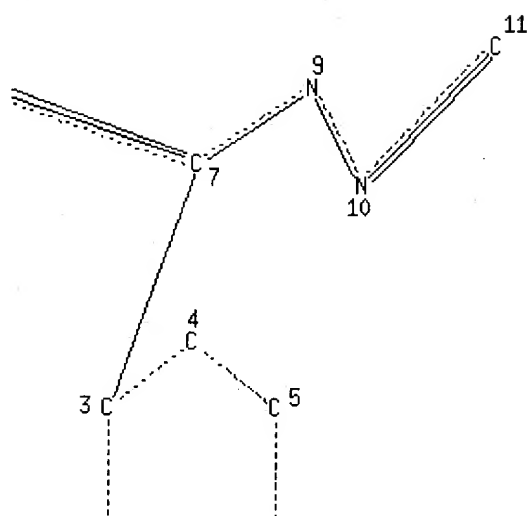
0 26 S 27

PRO

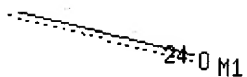
8 0



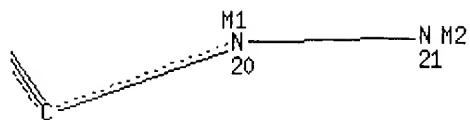
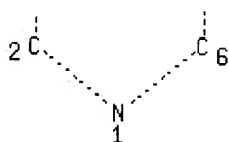
Page 1-A



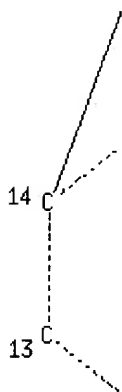
Page 1-B



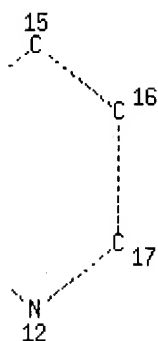
Page 2-A



Page 2-B



Page 3-A



Page 3-B

VAR G1=26/27

VAR G2=28/29

NODE ATTRIBUTES:

HCOUNT	IS	M1	AT	20
HCOUNT	IS	M2	AT	21
HCOUNT	IS	M1	AT	24
HCOUNT	IS	M3	AT	28
HCOUNT	IS	M2	AT	29
HCOUNT	IS	E3	AT	30
NSPEC	IS	R	AT	1
NSPEC	IS	R	AT	2
NSPEC	IS	R	AT	3
NSPEC	IS	R	AT	4
NSPEC	IS	R	AT	5
NSPEC	IS	R	AT	6
NSPEC	IS	C	AT	7
NSPEC	IS	C	AT	8
NSPEC	IS	C	AT	9
NSPEC	IS	C	AT	10
NSPEC	IS	RC	AT	11
NSPEC	IS	R	AT	12
NSPEC	IS	R	AT	13
NSPEC	IS	R	AT	14
NSPEC	IS	R	AT	15
NSPEC	IS	R	AT	16
NSPEC	IS	R	AT	17



NSPEC IS C AT 18
 NSPEC IS C AT 19
 NSPEC IS C AT 20
 NSPEC IS C AT 21
 NSPEC IS C AT 22
 NSPEC IS RC AT 23
 NSPEC IS C AT 24
 NSPEC IS C AT 25
 DEFAULT MLEVEL IS ATOM
 MLEVEL IS CLASS AT 7 8 9 10 11 18 19 20 21 23 24 26 27 28 29 30
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 30

STEREO ATTRIBUTES: NONE

=> s l16

SAMPLE SEARCH INITIATED 15:58:58 FILE 'CASREACT'

SCREENING COMPLETE - 33 REACTIONS TO VERIFY FROM 7 DOCUMENTS

100.0% DONE 33 VERIFIED 6 HIT RXNS 1 DOCS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 316 TO 1004

PROJECTED ANSWERS: 1 TO 79

L17 1 SEA SSS SAM L16 (6 REACTIONS)

=> s l16 full

THE ESTIMATED SEARCH COST FOR FILE 'CASREACT' IS 102.30 U.S. DOLLARS

DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 15:59:05 FILE 'CASREACT'

SCREENING COMPLETE - 419 REACTIONS TO VERIFY FROM 77 DOCUMENTS

100.0% DONE 419 VERIFIED 6 HIT RXNS 1 DOCS

SEARCH TIME: 00.00.01

L18 1 SEA SSS FUL L16 (6 REACTIONS)

=> d his

(FILE 'HOME' ENTERED AT 15:26:37 ON 09 JUL 2004)

FILE 'REGISTRY' ENTERED AT 15:26:43 ON 09 JUL 2004

E 310427-67-9/RN

L1 1 S E3

E 310427-65-7/RN

L2 1 S E3

E 10325-94-7/RN

L3 1 S E3

FILE 'CASREACT' ENTERED AT 15:40:30 ON 09 JUL 2004

L4 STRUCTURE UPLOADED

L5 11 S L4

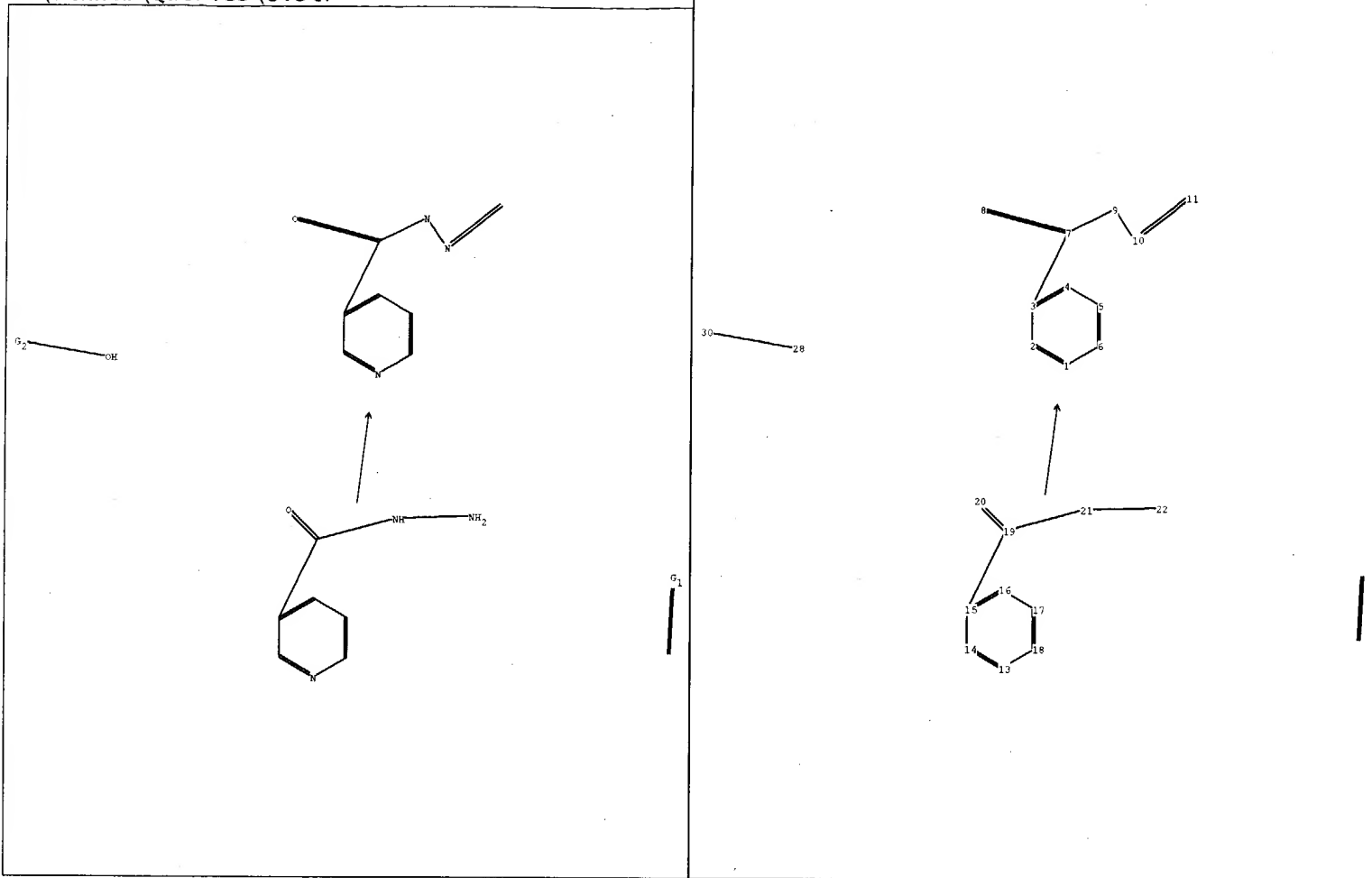
L6 187 S L4 FULL

L7 STRUCTURE UPLOADED
L8 10 S L7
L9 157 S L7 FULL

FILE 'REGISTRY' ENTERED AT 15:47:49 ON 09 JUL 2004
E ABSOLUTE ETHANOL/CN

FILE 'CASREACT' ENTERED AT 15:48:05 ON 09 JUL 2004
L10 STRUCTURE UPLOADED
L11 1 S L10
L12 3 S L11 FULL
L13 STRUCTURE UPLOADED
L14 1 S L13
L15 1 S L13 FULL
L16 STRUCTURE UPLOADED
L17 1 S L16
L18 1 S L16 FULL

=> s l18 not l15
L19 0 L18 NOT L15



chain nodes :

7 8 9 10 19 20 21 22 24 28 30

ring nodes :

1 2 3 4 5 6 13 14 15 16 17 18

ring/chain nodes :

11 25

chain bonds :

3-7 7-9 7-8 9-10 10-11 15-19 19-21 19-20 21-22 24-25 28-30

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18

exact/norm bonds :

7-9 7-8 9-10 10-11 19-21 19-20 24-25 28-30

exact bonds :

3-7 15-19 21-22

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18

isolated ring systems :

containing 1 : 13 :

G1:O,S

G2:CH3,Et

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS
 11:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:CLASS 20:CLASS
 21:CLASS 22:CLASS 24:CLASS 25:CLASS 28:CLASS 30:CLASS

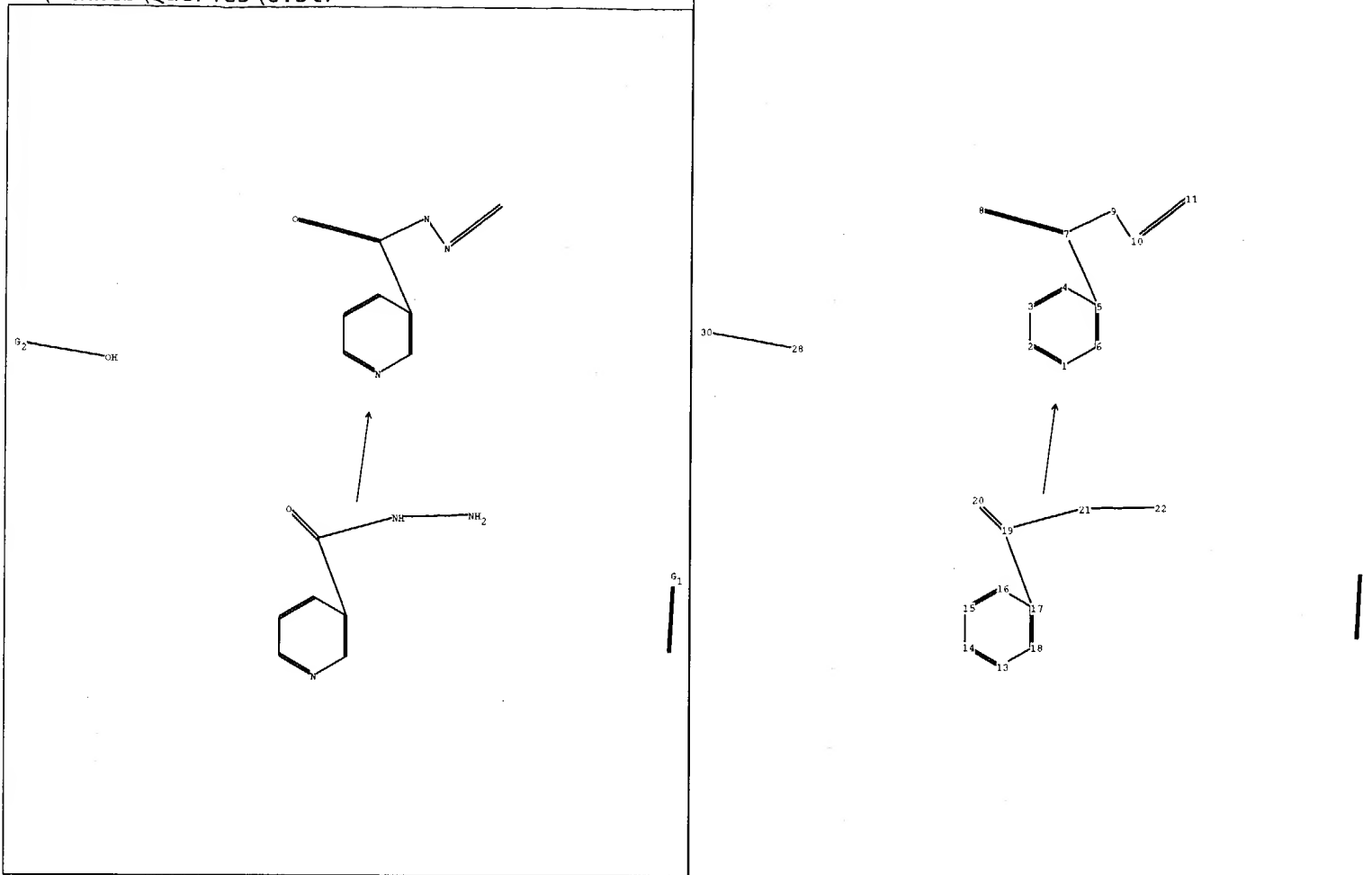
fragments assigned reactant role:

containing 13
 containing 24

fragments assigned product role:

containing 1

fragments assigned reactant/reagent role:



chain nodes :
 7 8 9 10 19 20 21 22 24 28 30
 ring nodes :
 1 2 3 4 5 6 13 14 15 16 17 18
 ring/chain nodes :
 11 25
 chain bonds :
 5-7 7-9 7-8 9-10 10-11 17-19 19-21 19-20 21-22 24-25 28-30
 ring bonds :
 1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18
 exact/norm bonds :
 7-9 7-8 9-10 10-11 19-21 19-20 24-25 28-30
 exact bonds :
 5-7 17-19 21-22
 normalized bonds :
 1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18
 isolated ring systems :
 containing 1 : 13 :

G1:O,S

G2:CH3,Et

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS
 11:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:CLASS 20:CLASS
 21:CLASS 22:CLASS 24:CLASS 25:CLASS 28:CLASS 30:CLASS

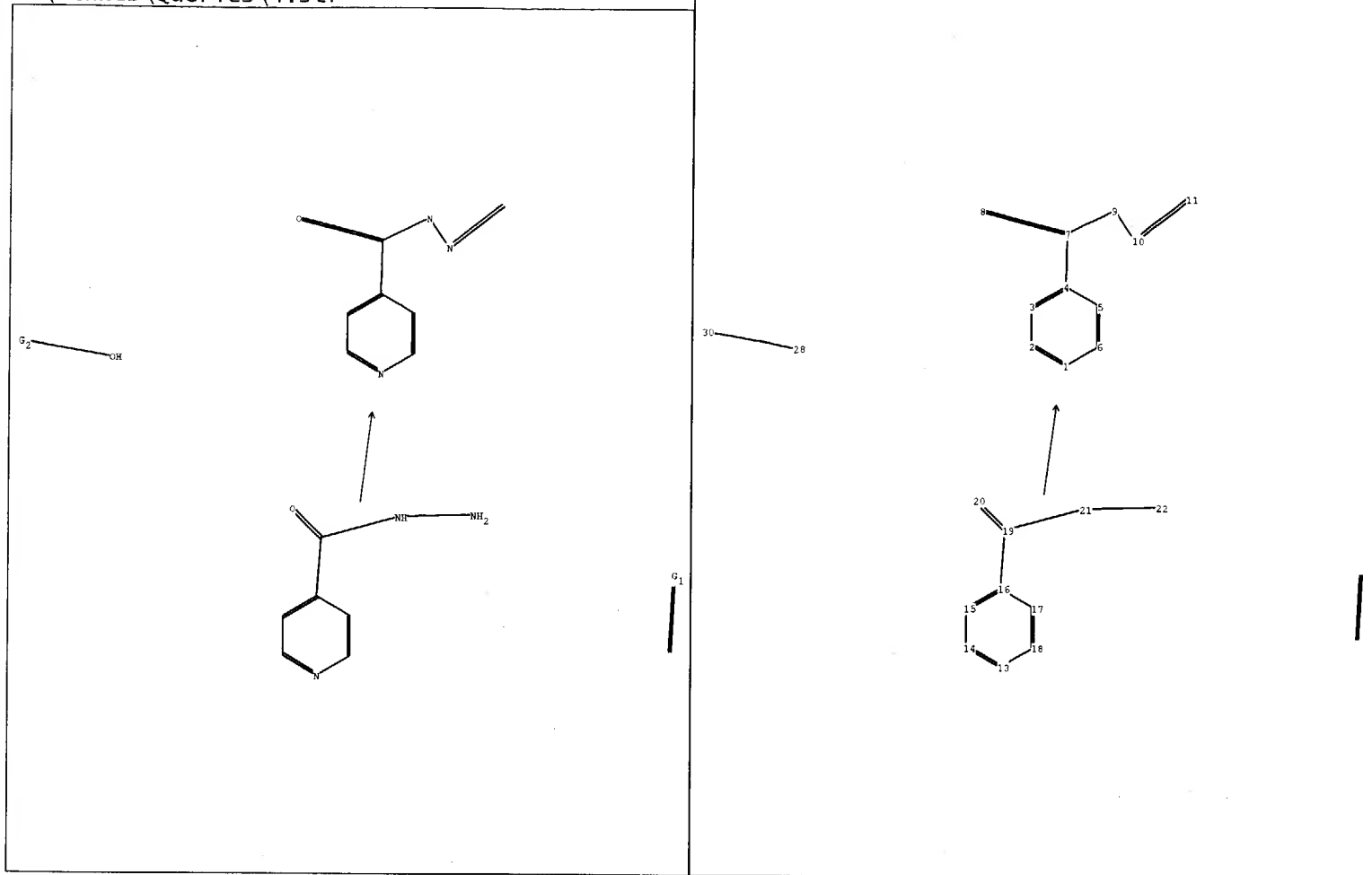
fragments assigned reactant role:

containing 13
 containing 24

fragments assigned product role:

containing 1

fragments assigned reactant/reagent role:



chain nodes :
 7 8 9 10 19 20 21 22 24 28 30
 ring nodes :
 1 2 3 4 5 6 13 14 15 16 17 18
 ring/chain nodes :
 11 25
 chain bonds :
 4-7 7-9 7-8 9-10 10-11 16-19 19-21 19-20 21-22 24-25 28-30
 ring bonds :
 1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18
 exact/norm bonds :
 7-9 7-8 9-10 10-11 19-21 19-20 24-25 28-30
 exact bonds :
 4-7 16-19 21-22
 normalized bonds :
 1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18
 isolated ring systems :
 containing 1 : 13 :

G1:O,S

G2:CH3,Et

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS
 11:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:CLASS 20:CLASS
 21:CLASS 22:CLASS 24:CLASS 25:CLASS 28:CLASS 30:CLASS

fragments assigned reactant role:

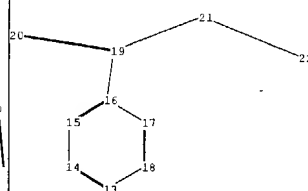
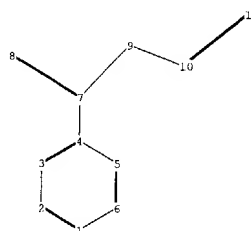
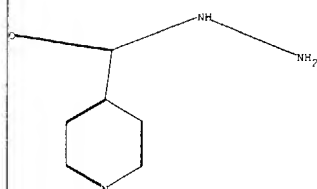
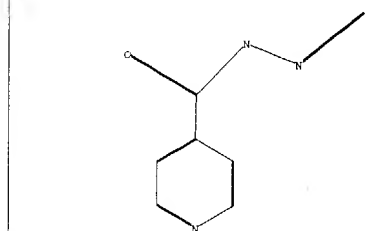
containing 13
 containing 24

fragments assigned product role:

containing 1

fragments assigned reactant/reagent role:

containing 28



chain nodes :

7 8 9 10 19 20 21 22 23 24 26

ring nodes :

1 2 3 4 5 6 13 14 15 16 17 18

ring/chain nodes :

11 25

chain bonds :

4-7 7-8 7-9 9-10 10-11 16-19 19-20 19-21 21-22 23-24 25-26

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18

exact/norm bonds :

7-8 7-9 9-10 10-11 19-20 19-21 23-24 25-26

exact bonds :

4-7 16-19 21-22

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18

isolated ring systems :

containing 1 : 13 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS
11:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:CLASS 20:CLASS
21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS

fragments assigned reactant role:

containing 13

fragments assigned product role:

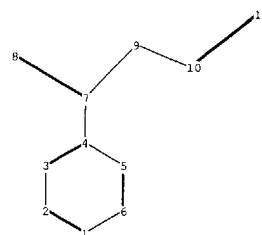
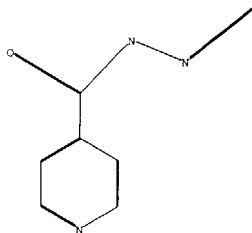
containing 1

fragments assigned reactant/reagent role:

containing 23

containing 25

C:\stnweb\Queries\3.str



chain nodes :

7 8 9 10

ring nodes :

1 2 3 4 5 6

ring/chain nodes :

11

chain bonds :

4-7 7-8 7-9 9-10 10-11

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6

exact/norm bonds :

7-8 7-9 9-10 10-11

exact bonds :

4-7

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6

isolated ring systems :

containing 1 :

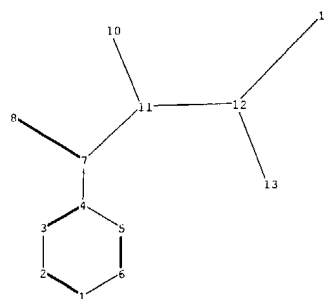
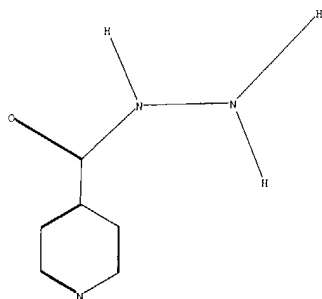
Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS

11:CLASS

fragments assigned product role:

containing 1



chain nodes :

7 8 10 11 12 13 14

ring nodes :

1 2 3 4 5 6

chain bonds :

4-7 7-8 7-11 10-11 11-12 12-13 12-14

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6

exact/norm bonds :

7-8 7-11 11-12

exact bonds :

4-7 10-11 12-13 12-14

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6

isolated ring systems :

containing 1 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 10:CLASS 11:CLASS
12:CLASS 13:CLASS 14:CLASS

fragments assigned product role:

containing 1